EAST Search History

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	17	f0f1 adj atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03
L2	11	f0f1-atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03
L3	123	H+-atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03
L4	214	H+ adj atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03
L5	230	L1 or L2 or L3 or L4	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03
L6	6	L5 and (tomita.in. or yokota.in.)	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03
S1	33711	method and biotinylat\$	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 11:06
S2	899	(method and biotinylat\$).clm.	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 11:10
S3	17	S2 and biotinylat\$.ti.	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:40
S4	17	f0f1 adj atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:41
S5	11	f0f1-atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:41

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EAST Search History

S6	123	H+-atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:41
S7	214	H+ adj atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:41
S8	230	S4 or S5 or S6 or S7	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03
S9	4	S8 and ammoniagenes	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:41
S10	31	S8 and (coryneform or corynebacterium)	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:44
S11	47	S8 and subtilis	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 12:44
S12	17	f0f1 adj atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 13:17
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S15	214	H+ adj atpase	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 13:17
S16	230	S12 or S13 or S14 or S15	US-PGPUB; USPAT; EPO; JPO; DERWENT	OR	OFF	2006/05/12 14:03

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Page 1 of 1 Entrez PubMed





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> > May 9 2006 14:13:00

IUBMB Enzyme Nomenclature

EC 3.6.3.14

Common name: H⁺-transporting two-sector ATPase

Reaction: ATP + $H_2O + H_{in}^+ = ADP + phosphate + H_{out}^+$

Glossary: F_0 the "o" refers to oligomycin. F_0 is incorrect.

Other names: ATP synthase; F_1 -ATPase; F_0 -ATPase; F_0 -ATPase; F_0 -ATPase; mitochondrial ATPase; coupling factors (F_0 , F_1 and F_0); chloroplast ATPase; bacterial F_0 -ATPase

Systematic name: ATP phosphohydrolase (H⁺-transporting)

Comments: A multisubunit non-phosphorylated ATPase that is involved in the transport of ions. Large enzymes of mitochondria, chloroplasts and bacteria with a membrane sector (F_0, V_0, A_0) and a cytoplasmic-compartment sector (F_1, V_1, A_1) . The F-type enzymes of the inner mitochondrial and thylakoid membranes act as ATP synthases. All of the enzymes included here operate in a rotational mode, where the extramembrane sector (containing 3 α - and 3 β -subunits) is connected via the δ -subunit to the membrane sector by several smaller subunits. Within this complex, the γ - and ϵ -subunits, as well as the 9-12 c subunits rotate by consecutive 120° angles and perform parts of ATP synthesis. This movement is driven by the H⁺ electrochemical potential gradient. The V-type (in vacuoles and clathrincoated vesicles) and A-type (archebacterial) enzymes have a similar structure but, under physiological conditions, they pump H⁺ rather than synthesize ATP.

Links to other databases: BRENDA, EXPASY, KEGG, ERGO, PDB, CAS registry number:

References:

- 1. Boyer, P.D. The binding change mechanism for ATP synthase some probabilities and possibilities. *Biochim. Biophys. Acta* 1140 (1993) 215-250. [Medline UI: 93112640]
- 2. Abrahams, J.P., Leslie, A.G.W., Lutter, R. and Walker, J.F. Structure at 2.8 Å resolution of F₁-ATPase from bovine heart mitochondria. *Nature* 375 (1994) 621-628. [Medline UI: 94344236]
- 3. Blair, A., Ngo, L., Park, J., Paulsen, I.T. and Saier, M.H., Jr. Phylogenetic analyses of the homologous transmembrane channel-forming proteins of the F_oF₁-ATPases of bacteria, chloroplasts and mitochondria. *Microbiology* 142 (1996) 17-32. [Medline UI: 96146047]
- 4. Noji, H., Yasuda, R., Yoshida, M. and Kinosita, K., Jr. Direct observation of the rotation of F₁-ATPase. *Nature* 386 (1997) 299-302. [Medline UI: 97222141]

[EC 3.6.3.14 created 1984 as EC 3.6.1.34, transferred 2000 to EC 3.6.3.14]

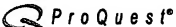
Return to EC 3.6.3 home page

EC 3.6.3.14 Page 2 of 2

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=>

(FILE 'HOME' ENTERED AT 14:04:27 ON 12 MAY 2006) FILE 'REGISTRY' ENTERED AT 14:05:15 ON 12 MAY 2006 L1 9 S 3.6.3.14 E "3.6.3.14"/CN 25 E "ATPASE"/CN 25 1 S E3 L2FILE 'MEDLINE, AGRICOLA, CAPLUS, BIOSIS, EMBASE, WPIDS' ENTERED AT 14:10:39 ON 12 MAY 2006 L3 3 S (L2 OR ATPASE) AND AMMONIAGENES L42 DUP REM L3 (1 DUPLICATE REMOVED) FILE 'CAPLUS' ENTERED AT 14:12:04 ON 12 MAY 2006 E TOMITA F/AU 25 L5 252 S (E18) E YOKOTA A/AU 25 L6 157 S (E29) L7 15 S (L5 OR L6) AND ATPASE L8 15 DUP REM L7 (0 DUPLICATES REMOVED) L9 15 S L8 L10 1 S L8 AND AMMONIAGENES



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14: gb_htg:* 15: gb_pl:*

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6	4131.2	43.5	8394	1	AB046112	AB046112 Corynebac
7	4130	43.5	8001	1	AB048368	AB048368 Brevibact

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From:

Steadman, David (AU1652)

Sent:

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NAME:

David Steadman

AU:

1656

Date:

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Remsen 2B05

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Please search the following sequence in commercial databases:

1) Standard search of SEQ ID NO:9 against nucleic acid databases.

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SEQ ID NO:9 against SEQ ID NO:21

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Searcher:	
Searcher Phone:	
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Date completed:	
Searcher Prep Time:	
Online Time:	

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S/L: O	ligomer:									
Encode/Trans	Encode/Transl:									
Structure #:_	Text:									
Inventor:	Litigation:									

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Title: US-10-694-779-21

Perfect score: 9500

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Scoring table: IDENTITY_NUC

Gapop 10.0 , Gapext 1.0

Searched: 1303057 seqs, 888780828 residues

Total number of hits satisfying chosen parameters: 2606114

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(without alignments)

12091.269 Million cell updates/sec

Title: US-10-694-779-21

Perfect score: 9500

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Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 8023312 seqs, 1165852854 residues

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Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications NA New:*

- 1: /cgn2 6/ptodata/1/pubpna/US08 NEW PUB.seq:*
- 2: /cgn2 6/ptodata/1/pubpna/US06 NEW PUB.seq:*
- 3: /cgn2_6/ptodata/1/pubpna/US07_NEW_PUB.seq:*
- 4: /cgn2_6/ptodata/1/pubpna/PCT_NEW_PUB.seq:*
- 5: /cgn2_6/ptodata/1/pubpna/US09 NEW PUB.seq:*
- 6: /cgn2 6/ptodata/1/pubpna/US09 NEW PUB.seq1:*
- 7: /cgn2_6/ptodata/1/pubpna/US10_NEW_PUB.seq:*
- 8: /cgn2_6/ptodata/1/pubpna/US10_NEW_PUB.seq1:*
- 9: /cgn2_6/ptodata/1/pubpna/US11_NEW_PUB.seq:*
- 10: /cgn2_6/ptodata/1/pubpna/US11 NEW PUB.seq2:*
- 11: /cgn2_6/ptodata/1/pubpna/US11 NEW PUB.seq3:*
- 12: /cgn2_6/ptodata/1/pubpna/US11_NEW_PUB.seq4:*
 13: /cgn2_6/ptodata/1/pubpna/US60_NEW_PUB.seq:*
- od No is the number of results andisted by shows to have

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		ક		
Result		Query		
No.	Score	Match Length	DB ID	Description
1	613.4	6.5 145761	.9 12 US-11-098-686-8739	Sequence 8739, Ap
2	508.8	5.4 1662	8 US-10-821-234-784	Sequence 784, App
3	508.2	5.3 3014	8 US-10-793-626-3680	Sequence 3680, Ap
4	494	5.2 1437	12 US-11-074-176-167	Sequence 167, App

```
5
        491.8
                  5.2
                        1413
                               8
                                  US-10-793-626-1813
                                                               Sequence 1813, Ap
    6
        485.6
                  5.1
                        1395
                               8
                                  US-10-467-657-1291
                                                               Sequence 1291, Ap
    7
                               7
                                                               Sequence 75392, A
        453.2
                  4.8
                        1638
                                  US-10-932-182A-75392
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        453.2
                  4.8
                        1638
                                  US-10-932-182A-75392
                                                               Sequence 75392, A
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                  4.7
                        1929
                               12
                                  US-11-119-351-9
                                                                Sequence 9, Appli
                  4.7
                        1647
                                  US-10-932-182A-2757
                                                               Sequence 2757, Ap
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        447.2
   11
        447.2
                  4.7
                        1647
                               7
                                  US-10-932-182A-2757
                                                               Sequence 2757, Ap
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                  4.7
                        1509
                                  US-11-074-176-349
                                                                Sequence 349, App
                               12
                  4.7
   13
        445.2
                        1518
                               12
                                  US-11-074-176-175
                                                                Sequence 175, App
   14
        438.2
                  4.6
                        1521
                               12
                                  US-11-098-686-9301
                                                                Sequence 9301, Ap
   15
        428.8
                  4.5
                        3080
                               8
                                  US-10-793-626-3855
                                                               Sequence 3855, Ap
          427
   16
                  4.5
                        1512
                               8
                                  US-10-793-626-1809
                                                               Sequence 1809, Ap
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                               7
                  4.5
                        1536
                                  US-10-932-182A-1477
                                                               Sequence 1477, Ap
   18
                  4.5
                               7
                                  US-10-932-182A-1477
        426.2
                        1536
                                                               Sequence 1477, Ap
   19
                  4.5
        425.6
                        1416
                               12
                                  US-11-098-686-9303
                                                                Sequence 9303, Ap
   20
        420.6
                  4.4
                        1545
                               8
                                  US-10-467-657-1287
                                                               Sequence 1287, Ap
   21
        406.6
                  4.3
                        1536
                                  US-10-932-182A-77398
                                                               Sequence 77398, A
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                  4.3
                        1536
                               7
                                  US-10-932-182A-77398
                                                               Sequence 77398, A
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        383.6
                  4.0 611587
                               12
                                  US-11-117-187-209
                                                                Sequence 209, App
С
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                               8
                                  US-10-793-626-4003
                                                               Sequence 4003, Ap
С
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                                  US-10-527-048-13
                                                               Sequence 13, Appl
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                        5834
                               7
                                  US-10-527-048-15
                                                               Sequence 15, Appl
С
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                        6465
                               7
                                  US-10-527-048-25
                                                               Sequence 25, Appl
C
C
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                        6659
                               7
                                  US-10-527-048-22
                                                               Sequence 22, Appl
  29
        340.8
                  3.6
                        7057
                               7
С
                                  US-10-527-048-26
                                                               Sequence 26, Appl
                               7
                                                               Sequence 19, Appl
С
   30
        340.8
                  3.6
                        7510
                                  US-10-527-048-19
                               7
   31
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                  3.6
                        7549
                                  US-10-527-048-24
                                                               Sequence 24, Appl
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                  3.6
                        8327
                               7
                                  US-10-527-048-23
                                                               Sequence 23, Appl
   33
        337.6
                  3.6
                        1887
                                  US-10-527-048-8
                                                               Sequence 8, Appli
  34
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                  3.5
                        1352
                               8
                                  US-10-485-517-72
                                                               Sequence 72, Appl
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                        1066
                                  US-11-136-527-4053
                               12
                                                                Sequence 4053, Ap
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                  2.4
                         902
                               12
                                   US-11-128-061-217
                                                                Sequence 217, App
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        225.6
                  2.4
                         902
                               12
                                   US-11-128-049-217
                                                                Sequence 217, App
   38
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                  2.2
                         600
                               12
                                   US-11-128-061-3859
                                                                Sequence 3859, Ap
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                               12
        211.4
                                  US-11-128-049-3859
                                                                Sequence 3859, Ap
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                         517
                                  US-09-925-065A-327316
                               6
                                                               Sequence 327316,
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                         517
                               6
                                  US-09-925-065A-327317
                                                               Sequence 327317,
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С
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                               6
                                  US-09-925-065A-611756
                                                               Sequence 611756,
С
   43
        204.6
                  2.2
                         573
                               6
                                  US-09-925-065A-611757
                                                               Sequence 611757,
С
   44
        197.4
                  2.1
                         542
                               6
                                  US-09-925-065A-327793
                                                               Sequence 327793,
   45
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                         542
                               6
                  2.1
                                  US-09-925-065A-327794
                                                               Sequence 327794,
```

OM nucleic - nucleic search, using sw model

March 15, 2006, 05:44:57 ; Search time 1112 Seconds Run on:

(without alignments)

5466.007 Million cell updates/sec

Title: US-10-694-779-9

Perfect score: 912

Sequence: 1 atgtgcgacggagtccgtag.....cacttcacgcagactcgcac 912

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

4996997 seqs, 3332346308 residues Searched:

Total number of hits satisfying chosen parameters: 9993994

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : N Geneseq 21:*

Q.

1: geneseqn1980s:*

2: geneseqn1990s:*

3: geneseqn2000s:*

4: geneseqn2001as:*

5: geneseqn2001bs:*

6: geneseqn2002as:*

7: geneseqn2002bs:*

8: geneseqn2003as:*

9: geneseqn2003bs:*

10: geneseqn2003cs:*

11: geneseqn2003ds:*

12: geneseqn2004as:* 13: geneseqn2004bs:*

14: geneseqn2005s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	Query Match	Length	DB	ID	Description
1	912	100.0	912	6	ABL54659	Abl54659 Corynebac
2	912	100.0	9500	6	ABL54671	Abl54671 Corynebac
3	504.2	55.3	349980	5	AAH68528	Aah68528 C glutami

							-
	4	488.2	53.5	933	4	AAF71737	Aaf71737 Corynebac
	5	456.2	50.0	862	4	AAF71738	Aaf71738 Corynebac
	6	454	49.8	940	8	ACA29621	Aca29621 Prokaryot
	7	443.6	48.6	810	5	AAH66301	Aah66301 C glutami
С	8	118.8		349980	6	ABQ81844	Abq81844 Bifidobac
	9	50.8	5.6	843	14	ACL66536	Acl66536 M. xanthu
С	10	50.8	5.6	1394	14	ACL63902	Acl63902 M. xanthu
	11	48.8	5.4	4364	4	AAS59660	Aas59660 Propionib
	12	48.8	5.4	4364	8	ACF64589	Acf64589 Propionib
	13	48.4	5.3	543	13	ADS59769	Ads59769 Bacterial
	14	48.4	5.3	732	13	ADS62938	Ads62938 Bacterial
	15	48.4	5.3	732	13	ADS62291	Ads62291 Bacterial
	16	48.4	5.3	732	13	ADS62785	Ads62785 Bacterial
	17	48.4	5.3	849	8	ACA25438	Aca25438 Prokaryot
	18	47.2	5.2	849	8	ACA26013	Aca26013 Prokaryot
С	19	47.2	5.2	2000	8	ADA71938	Ada71938 Rice gene
	20	44.6	4.9	1703	4	AAD02475	Aad02475 cDNA clon
	21	43.8	4.8	2000	8	ADA71938	Ada71938 Rice gene
	22	43	4.7	879	8	ACA27392	Aca27392 Prokaryot
	23	40.2	4.4	849	8	ACA23312	Aca23312 Prokaryot
С	24	40	4.4	680	3	AAF08703	Aaf08703 Fusarium
C	25	40	4.4	680	13	ADU52744	Adu52744 Fusarium
С	26	40	4.4	680	14	ADZ90747	Adz90747 Fusarium
С	27	39	4.3	1581	3	AAA93691	Aaa93691 Agrobacte
С	28	39	4.3	1581	4	AAF86183	Aaf86183 Mannose i
C	29	39	4.3	1581	8	ACA61518	Aca61518 Modified
	30	38	4.2	693	8	ACA25636	Aca25636 Prokaryot
	31	37.8	4.1	702	14	ACL70731	Acl70731 M. xanthu
	32	37.8	4.1	726	13	ADT44413	Adt44413 Bacterial
С	33	37.8	4.1	6430	14	ACL64206	Acl64206 M. xanthu
	34	37	4.1	663	11	ACH94558	Ach94558 Klebsiell
	35	37	4.1	696	8	ACA25755	Aca25755 Prokaryot
	36	36.6	4.0	795	14	ADY77504	Ady77504 A. thalia
	37	36.6	4.0	1813	12	ADM47591	Adm47591 Polynucle
	38	36.6	4.0	1865	4	AAD02473	Aad02473 cDNA clon
С	39	36.6	4.0	35829	4	AAS59573	Aas59573 Propionib
С	40	36.6	4.0	35829	8	ACF64502	Acf64502 Propionib
	41	36.4	4.0	1256	13	ADX35007	Adx35007 Plant ful
	42	36.4	4.0	2145	9	ADA48625	Ada48625 Rice gene
	43	36.4	4.0	2145	11	ACL30288	Acl30288 Rice abio
	44	36.4	4.0	2187	8	ADA69492	Ada69492 Rice gene
	45	36.4	4.0	2189	9	ADA48499	Ada48499 Rice gene

OM nucleic - nucleic search, using sw model

Run on: March 15, 2006, 22:16:24; Search time 1455 Seconds

(without alignments)

5183.280 Million cell updates/sec

Title: US-10-694-779-9

Perfect score: 912

Sequence: 1 atgtgcgacggagtccgtag.....cacttcacgcagactcgcac 912

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 9793542 seqs, 4134689005 residues

Total number of hits satisfying chosen parameters: 19587084

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications NA Main:*

- 1: /cgn2 6/ptodata/1/pubpna/US07 PUBCOMB.seq:*
- 2: /cgn2 6/ptodata/1/pubpna/US08 PUBCOMB.seq:*
- 3: /cgn2_6/ptodata/1/pubpna/US09A_PUBCOMB.seq:*
- 4: /cgn2_6/ptodata/1/pubpna/US09B_PUBCOMB.seq:*
- 5: /cgn2_6/ptodata/1/pubpna/US10A_PUBCOMB.seq:*
- 6: /cgn2_6/ptodata/1/pubpna/US10B_PUBCOMB.seq:*
- 7: /cgn2_6/ptodata/1/pubpna/US10C_PUBCOMB.seq:*
- 8: /cgn2_6/ptodata/1/pubpna/US10D_PUBCOMB.seq:*
- 9: /cgn2_6/ptodata/1/pubpna/US10E_PUBCOMB.seq:*
- 10: /cgn2_6/ptodata/1/pubpna/US11_PUBCOMB.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		8				
Result		Query				
No.	Score	Match	Length 1	OB	ID	Description
1	912	100.0	912	3	US-09-901-884-9	Sequence 9, Appli
2	912	100.0	912	8	US-10-694-779-9	Sequence 9, Appli
3	912	100.0	9500	3	US-09-901-884-21	Sequence 21, Appl
4	912	100.0	9500	8	US-10-694-779-21	Sequence 21, Appl
5	504.2	55.3	3309400	3	US-09-738-626-1	Sequence 1, Appli
6	488.2	53.5	933	8	US-10-781-014-755	Sequence 755, App
7	456.2	50.0	862	8	US-10-781-014-757	Sequence 757, App

```
454
                49.8
                         940
                                 US-10-282-122A-17491
                                                             Sequence 17491, A
    8
        443.6
                         810
                              3
                                 US-09-738-626-1336
                                                             Sequence 1336, Ap
    9
                48.6
                                                             Sequence 2874, Ap
                         807
                              6
                                 US-10-156-761-2874
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        119.4
                13.1
   11
        119.4
                13.1 125746
                                 US-10-156-761-15102
                                                             Sequence 15102, A
                13.1 9025608
   12
        119.4
                              6
                                 US-10-156-761-1
                                                              Sequence 1, Appli
C
                13.0 2256646 7
                                  US-10-470-565-1
                                                              Sequence 1, Appli
С
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        118.8
         48.4
                 5.3
                         543
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                                                             Sequence 35443, A
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         48.4
                 5.3
                         732
                              6
                                 US-10-369-493-37965
                                                             Sequence 37965, A
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                              6
                                 US-10-369-493-38459
                                                             Sequence 38459, A
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                         732
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         48.4
                 5.3
                             6
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                                                             Sequence 38612, A
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                                 US-10-282-122A-13308
                                                             Sequence 13308, A
                              7
                                                             Sequence 13883, A
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                 5.2
                         849
                                 US-10-282-122A-13883
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                         864
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                                 US-10-425-115-56270
                                                             Sequence 56270, A
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                              7
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                                 US-10-282-122A-15262
                                                             Sequence 15262, A
                        1160
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         42.6
                 4.7
                             5
                                 US-10-123-155-234
                                                             Sequence 234, App
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         42.6
                 4.7
                        1160
                                 US-10-146-731-234
                                                             Sequence 234, App
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                 4.7
                        1160
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                                 US-10-140-472-234
                                                             Sequence 234, App
   25
         42.6
                 4.7
                        1160
                             6
                                 US-10-141-761-234
                                                             Sequence 234, App
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         42.6
                 4.7
                        1160
                             6
                                 US-10-142-885-234
                                                             Sequence 234, App
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         42.6
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                        1160
                             6
                                 US-10-158-790-234
                                                             Sequence 234, App
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                 4.7
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                                 US-10-137-871-234
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                        1160 6
                                 US-10-140-923-234
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                                                             Sequence 234, App
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                 4.7
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                                 US-10-141-759-234
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                                 US-10-140-805-234
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         42.6
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                                 US-10-140-864-234
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С
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                                 US-10-437-963-1603
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                                 US-10-282-122A-13625
                                                             Sequence 13625, A
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                 4.0
                         594 5
С
                                 US-10-123-155-10
                                                             Sequence 10, Appl
   45
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                         594 6
                                 US-10-146-731-10
C
                 4.0
                                                             Sequence 10, Appl
```

OM nucleic - nucleic search, using sw model

Run on: March 15, 2006, 22:58:26; Search time 3731 Seconds

(without alignments)

11436.564 Million cell updates/sec

Title: US-10-694-779-9

Perfect score: 912

Sequence: 1 atgtgcgacggagtccgtag.....cacttcacgcagactcgcac 912

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 41078325 seqs, 23393541228 residues

Total number of hits satisfying chosen parameters: 82156650

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : EST:*

1: gb_est1:*

2: gb_est2:*

3: gb_est3:*

4: gb_htc:*

5: gb_est4:*

6: gb est5:*

7: gb_est6:*

8: gb_est7:*

9: gb_gss1:*

10: gb_gss2:*

11: gb_gss3:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

D = -	T &-		%				
kes	sult No.	Score	Query Match	Length	DB	ID	Description
С	1	81.6	8.9	426	2	BI062358	BI062358 IL3-UT011
	2	47.6	5.2	1069	8	DR738787	DR738787 FGAS08400
	3	46.6	5.1	839	10	AG882978	AG882978 Oryza sat
	4	46	5.0	475	8	DN180328	DN180328 HO27G01S
	5	46	5.0	612	8	DN181015	DN181015 HO25H19S
	6	46	5.0	1000	1	AU252302	AU252302 AU252302

	7	46	5.0	1316	1	AU252292	AU252292 AU252292
	8	45.6	5.0	654	10	CW138494	CW138494 104_528_1
	9	44.6	4.9	405	8	DN180329	DN180329 HO27G02S
	10	44.6	4.9	835	8	CV776873	CV776873 FGAS07127
	11	44.6	4.9	1146	7	CK166509	CK166509 FGAS05066
	12	43.6	4.8	687	10	AG946800	AG946800 Drosophil
	13	43	4.7	662	3	BJ281216	BJ281216 BJ281216
	14	43	4.7	695	6	CD882228	CD882228 F1.105M13
	15	43	4.7	857	8	CV763444	CV763444 FGAS05783
	16	43	4.7	969	8	CV770252	CV770252 FGAS06464
	17	43	4.7	1105	8	DR739375	DR739375 FGAS08459
	18	43	4.7	1169	7	CK162000	CK162000 FGAS01458
	19	42.6	4.7	257	1	AL828709	AL828709 AL828709
	20	42.6	4.7	443	6	CD461012	CD461012 Fg09_03g1
C	21	42	4.6	501	6	CD116784	CD116784 ME1-0043T
C	22	42	4.6	593	10	AG937790	AG937790 Drosophil
C	23	41.4	4.5	577	5	BU060496	BU060496 Fgr-C_1_D
	24	41.4	4.5	581	5	BU060034	BU060034 Fgr-C_0_J
C	25	41.4	4.5	609	5	BU063571	BU063571 Fgr_3_E16
C	26	41.4	4.5	635	5	BU065578	BU065578 Fgr_7_P05
	27	41.4	4.5	661	5	BU062199	BU062199 Fgr_1_H15
	28	41.4	4.5	669	5	BU060497	BU060497 Fgr-C_1_D
	29	41.4	4.5	670	5	BU063126	BU063126 Fgr_2_L07
	30	41.4	4.5	704	5	BU061251	BU061251 FgrN_5_M1
	31	41.4	4.5	712	5	BU063572	BU063572 Fgr_3_E16
	32	40.4	4.4	823	7	CK200509	CK200509 FGAS00902
	33	40.2	4.4	997	10	CNS005TE	AL060767 Drosophil
C	34	39.8	4.4	488	5	BU063125	BU063125 Fgr_2_L07
C	35	39.8	4.4	494	5	BU060033	BU060033 Fgr-C_0_J
	36	39	4.3	568	5	BQ838054	BQ838054 WHE2906_A
	37	39	4.3	636	3	BI954020	BI954020 HVSMEm001
	38	38.8	4.3	530	3	BJ244886	BJ244886 BJ244886
	39	38.8	4.3	562	3	BI774730	BI774730 466936 MA
С	40	38.8	4.3	741	7	CK358958	CK358958 AGENCOURT
	41	38.8	4.3	861	7	CK203625	CK203625 FGAS01215
С	42	38.4	4.2	458	7	CF964603	CF964603 12818rsic
С	43	38.4	4.2	573	10	CZ632396	CZ632396 OM Ba017
	44	38.4	4.2	779	7	CO387234	CO387234 AGENCOURT
C	45	38.2	4.2	470	5	BU062198	BU062198 Fgr 1 H15
							y

OM nucleic - nucleic search, using sw model

Run on: March 17, 2006, 09:15:55; Search time 4968 Seconds

(without alignments)

12744.481 Million cell updates/sec

Title: US-10-694-779-21

Perfect score: 9500

Sequence: 1 acctcgtatgggcagtaatt.....acgggcaacgcgtccgcggg 9500

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 4996997 seqs, 3332346308 residues

Total number of hits satisfying chosen parameters: 99933994

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : N Geneseq 21:*

1: geneseqn1980s:*

2: geneseqn1990s:*

3: geneseqn2000s:*

4: geneseqn2001as:*

5: geneseqn2001bs:*

6: geneseqn2002as:*

7: geneseqn2002bs:*

8: geneseqn2003as:*

9: geneseqn2003bs:*

10: geneseqn2003cs:*

10. genesequizouses.

11: geneseqn2003ds:*
12: geneseqn2004as:*

13: geneseqn2004bs:*

14: geneseqn2005s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

		*					
Result		Query					
No.	Score	Match	Length	DB	ID		Description
1	9500	100.0	9500	6	ABL54671		Abl54671 Corynebac
2	4227.4	44.5	349980	5	AAH68528		Aah68528 C glutami
3	1749.2	18.4	110000	4	AA199682	14	Continuation (15 o

	4	1749.2		110000	4	AAI99683_14	Continuation (15 o
C	5	1732.4	18.2	36241		ADB74385	Adb74385 Mycobacte
С	6	1667.2		349980	6	ABQ81844	Abq81844 Bifidobac
	7	1638	17.2	1638	6	ABL54663	Abl54663 Corynebac
С	8	1474.2	15.5	20798	4	AAS59620	Aas59620 Propionib
С	9	1474.2	15.5	20798	8	ACF64549	Acf64549 Propionib
	10	1443	15.2	1443	6	ABL54665	Abl54665 Corynebac
	11	1170.8	12.3	1572	4	AAF71740	Aaf71740 Corynebac
	12	1159	12.2	1764	4	AAF71739	Aaf71739 Corynebac
	13	1158.6	12.2	1674	5	AAH66305	Aah66305 C glutami
	14	1137.2	12.0	1659	8	ACA29625	Aca29625 Prokaryot
	15	1097.8	11.6	1452	2	AAT03954	Aat03954 Brevibact
	16	1092.6	11.5	1449	5	AAH66307	Aah66307 C glutami
	17	981	10.3	981	6	ABL54664	Abl54664 Corynebac
	18	912	9.6	912	6	ABL54659	Abl54659 Corynebac
	19	853.8	9.0	8008	2	AAZ32024	Aaz32024 Human MET
	20	853.8	9.0	8009	5	AAC90081	Aac90081 X99599 cD
	21	849.8	8.9	110000	6	ABQ67196_5	Continuation (6 of
C	22	849.8		110000	6	ABQ69245_26	Continuation (27 o
C	23	833.8	8.8	110000	6	ABA03041_25	Continuation (26 o
С	24	833.8	8.8	110000	6	ABA03041_26	Continuation (27 o
	25	813	8.6	813	6	ABL54662	Abl54662 Corynebac
	26	805	8.5	1662	8	ACA38144	Aca38144 Prokaryot
	27	799.4	8.4	1920	5	AAS03789	Aas03789 Mycobacte
	28	780.4	8.2	10399	2	AAV52293	Aav52293 Streptoco
C	29	780.4	8.2	110000	10	ABS56454_13	Continuation (14 o
	30	772	8.1	1521	13	ADS56270	Ads56270 Bacterial
C	31	769.2	8.1	46593	3	AAA81456	Aaa81456 N. mening
C	32	769.2		110000	3	AAA81489_5	Continuation (6 of
C	33	769.2		349980	3	AAF21612	Aaf21612 Neisseria
	34	765.2	8.1	1647	8	ACA38774	Aca38774 Prokaryot
	35	765.2	8.1	1650	8	ACA40479	Aca40479 Prokaryot
	36	758.6	8.0	4815	2	AAV29571	Aav29571 L. lactis
	37	757.4	8.0	1413		ADS56268	Ads56268 Bacterial
	38	751.4	7.9	7750	2	AAX13153	Aax13153 Enterococ
	39	751.4	7.9	7750	6	ABS98948	Abs98948 Enterococ
	40	751		110000	10	ADF77343_10	Continuation (11 o
	41	734.2	7.7	71979	13	ADV87736	Adv87736 Streptoco
	42	734.2	7.7	71979	13	ADV78989	Adv78989 Streptoco
	43	734.2		110000	6	ABN71527_08	Continuation (9 of
	44	734.2		110000	13	ADV81204_09	Continuation (10 o
	45	732.8	7.7	1677	8	ACA39764	Aca39764 Prokaryot

OM nucleic - nucleic search, using sw model

Run on: March 17, 2006, 19:30:20; Search time 4417 Seconds

(without alignments)

17785.622 Million cell updates/sec

Title: US-10-694-779-21

Perfect score: 9500

Sequence: 1 acctcgtatgggcagtaatt.....acgggcaacgcgtccgcggg 9500

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 9793542 seqs, 4134689005 residues

Total number of hits satisfying chosen parameters: 19587084

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

٩.

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications NA Main:*

- 1: /cgn2 6/ptodata/1/pubpna/US07 PUBCOMB.seq:*
- 2: /cgn2_6/ptodata/1/pubpna/US08_PUBCOMB.seq:*
- 3: /cgn2 6/ptodata/1/pubpna/US09A PUBCOMB.seq:*
- 4: /cgn2_6/ptodata/1/pubpna/US09B_PUBCOMB.seq:*
- 5: /cgn2_6/ptodata/1/pubpna/US10A_PUBCOMB.seq:*
- 6: /cgn2 6/ptodata/1/pubpna/US10A_PUBCOMB.seq:*
- 7: /cgn2 6/ptodata/1/pubpna/US10C PUBCOMB.seq:*
- 8: /cgn2 6/ptodata/1/pubpna/US10D PUBCOMB.seq:*
- 9: /cgn2 6/ptodata/1/pubpna/US10E PUBCOMB.seq:*
- 10: /cgn2_6/ptodata/1/pubpna/US11_PUBCOMB.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

	_		70				
Res	ult		Query				
	No.	Score	Match	Length 1	DΒ	ID	Description
	1	9500	100.0	9500	3	US-09-901-884-21	Sequence 21, Appl
	2	9500	100.0	9500	8	US-10-694-779-21	Sequence 21, Appl
	3	4227.4	44.5	3309400	3	US-09-738-626-1	Sequence 1, Appli
C	4	1667.2	17.5	2256646	7	US-10-470-565-1	Sequence 1, Appli
	5	1665.2	17.5	125746	6	US-10-156-761-15102	Sequence 15102, A
С	6	1665.2	17.5	9025608	6	US-10-156-761-1	Sequence 1, Appli
	7	1638	17.2	1638	3	US-09-901-884-13	Sequence 13, Appl

```
17.2
                        1638
                                 US-10-694-779-13
                                                             Sequence 13, Appl
    8
         1638
                              8
                                                             Sequence 15, Appl
    9
         1443
                15.2
                        1443
                              3
                                 US-09-901-884-15
                                                             Sequence 15, Appl
   10
         1443
                15.2
                        1443
                              8
                                 US-10-694-779-15
   11
       1170.8
                12.3
                        1572
                              8
                                 US-10-781-014-761
                                                             Sequence 761, App
                12.2
                        1764
                                 US-10-781-014-759
                                                             Sequence 759, App
   12
         1159
                              8
   13
       1158.6
                12.2
                        1674
                              3
                                 US-09-738-626-1340
                                                             Sequence 1340, Ap
       1137.2
                12.0
                        1659
                                 US-10-282-122A-17495
                                                             Sequence 17495, A
   14
   15
       1092.6
                11.5
                        1449
                              3
                                 US-09-738-626-1342
                                                             Sequence 1342, Ap
          981
                10.3
                         981
                              3
                                 US-09-901-884-14
                                                             Sequence 14, Appl
   16
   17
          981
                10.3
                         981
                              8
                                 US-10-694-779-14
                                                             Sequence 14, Appl
   18
          912
                 9.6
                         912
                              3
                                 US-09-901-884-9
                                                             Sequence 9, Appli
                 9.6
   19
          912
                         912
                              8
                                 US-10-694-779-9
                                                             Sequence 9, Appli
   20
        853.8
                 9.0
                        8009
                              3
                                 US-09-373-658-34
                                                             Sequence 34, Appl
   21
        853.8
                 9.0
                        8009
                              3
                                 US-09-989-687-34
                                                             Sequence 34, Appl
   22
                 8.9 684707
                              7
                                 US-10-398-221-9
                                                             Sequence 9, Appli
        849.8
   23
        849.8
                 8.9 3011208
                                 US-10-398-221-2058
                                                              Sequence 2058, Ap
   24
          813
                 8.6
                         813
                              3
                                 US-09-901-884-12
                                                             Sequence 12, Appl
   25
          813
                 8.6
                         813
                                 US-10-694-779-12
                                                             Sequence 12, Appl
                              8
   26
          805
                 8.5
                        1662
                              7
                                 US-10-282-122A-26014
                                                             Sequence 26014, A
   27
        780.4
                 8.2
                      10399
                              2
                                 US-08-961-527-160
                                                             Sequence 160, App
   28
        780.4
                 8.2
                      10399
                              7
                                 US-10-158-844-160
                                                             Sequence 160, App
   29
        780.4
                 8.2 2162598 8
                                 US-10-472-928-4979
                                                              Sequence 4979, Ap
   30
          772
                 8.1
                        1521
                              6
                                 US-10-369-493-31944
                                                             Sequence 31944, A
        769.2
С
   31
                      46594
                                 US-10-915-740A-4
                 8.1
                              9
                                                             Sequence 4, Appli
        769.2
                 8.1 2242716 9
   32
                                 US-10-915-740A-1068
                                                              Sequence 1068, Ap
   33
        765.2
                        1647
                 8.1
                              7
                                 US-10-282-122A-26644
                                                             Sequence 26644, A
   34
        765.2
                              7
                 8.1
                        1650
                                 US-10-282-122A-28349
                                                             Sequence 28349, A
   35
        758.6
                 8.0
                        4815
                              6
                                 US-10-310-630-1
                                                             Sequence 1, Appli
   36
        757.4
                 8.0
                        1413
                              6
                                 US-10-369-493-31942
                                                             Sequence 31942, A
   37
        751.4
                 7.9
                        7750
                                                             Sequence 216, App
                              3
                                 US-09-070-927A-216
                 7.8
   38
        739.2
                        1587
                              6
                                 US-10-156-761-2870
                                                             Sequence 2870, Ap
                                                             Sequence 27634, A
   39
        732.8
                 7.7
                        1677
                              7
                                 US-10-282-122A-27634
С
   40
          720
                 7.6 2731748
                              7
                                 US-10-297-465A-1
                                                              Sequence 1, Appli
   41
        708.8
                 7.5
                                 US-10-156-761-2868
                        1434
                             6
                                                             Sequence 2868, Ap
   42
        703.8
                 7.4
                        6901
                              8
                                 US-10-857-625-130
                                                             Sequence 130, App
   43
          629
                 6.6
                         778
                              8
                                 US-10-781-014-763
                                                             Sequence 763, App
   44
        603.8
                                                             Sequence 35121, A
                 6.4
                        1515
                              6
                                 US-10-369-493-35121
   45
        594.6
                 6.3
                        1527 6
                                 US-10-369-493-34379
                                                             Sequence 34379, A
```

OM nucleic - nucleic search, using sw model

Run on: March 17, 2006, 12:34:04; Search time 22063 Seconds

(without alignments)

20145.823 Million cell updates/sec

Title: US-10-694-779-21

Perfect score: 9500

Sequence: 1 acctcgtatgggcagtaatt.....acgggcaacgcgtccgcggg 9500

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 41078325 seqs, 23393541228 residues

Total number of hits satisfying chosen parameters: 82156650

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : EST:*

1: gb_est1:*

2: gb_est2:*

3: gb_est3:*

4: gb_htc:*

5: gb_est4:*

6: gb est5:*

7: gb_est6:*

8: gb_est7:*

9: gb_gss1:*

10: gb_gss2:*

11: gb gss3:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result		% Query				
No.	Score	Match	Length	DB	ID	Description
1	538.6	5.7	1613	4	CNSOEQFJ	CR636937 Tetraodon
2	529.6	5.6	2076	4	AY849656	AY849656 Magnaport
3	529.4	5.6	2196	4	AK043976	AK043976 Mus muscu
4	529.4	5.6	2197	4	AK076572	AK076572 Mus muscu
5	527.2	5.5	1821	4	CNS0EPRY	CR636088 Tetraodon
6	526.2	5.5	1847	4	BC020417	BC020417 Mus muscu

7	513	5.4	1910	4	CR861028	CR861028 Pongo pyg
8	510.8	5.4	1845	7	CN210866	CN210866 WLA073D01
9	510.8	5.4	1864	7	CN210865	CN210865 RJA108H07
10	510.4	5.4	1844	4	BC042694	BC042694 Homo sapi
11	509.8	5.4	1675	4	CNS0GQK6	CR730449 Tetraodon
12	508.8	5.4	1767	4	CR604755	CR604755 full-leng
13	508.8	5.4	1768	4	CR591462	CR591462 full-leng
14	508.8	5.4	1769	4	CR617088	CR617088 full-leng
15	508.8	5.4	1793	4	CR625409	CR625409 full-leng
16	508.8	5.4	1803	4	CR607960	CR607960 full-leng
17	508.8	5.4	1804	4	CR611390	CR611390 full-leng
18	508.8	5.4	1806	4	CR593595	CR593595 full-leng
19	508.8	5.4	1809	4	CR590601	CR590601 full-leng
20	508.8	5.4	1811	4	CR609973	CR609973 full-leng
21	508.8	5.4	1812	4	CR613857	CR613857 full-leng
22	508.8	5.4	1814	4	CR604007	CR604007 full-leng
23	508.8	5.4	1814	4	CR617331	CR617331 full-leng
24	508.8	5.4	1815	4	CR591138	CR591138 full-leng
25	508.8	5.4	1815	4	CR594700	CR594700 full-leng
26	508.8	5.4	1816	4	CR608051	CR608051 full-leng
27	508.8	5.4	1823	4	CR609069	CR609069 full-leng
28	508.8	5.4	1825	4	CR610155	CR610155 full-leng
29	508.8	5.4	1826	4	CR590562	CR590562 full-leng
30	508.8	5.4	1827	4	CR609904	CR609904 full-leng
31	508.8	5.4	1827	4	CR617880	CR617880 full-leng
32	508.8	5.4	1827	4	CR626813	CR626813 full-leng
33	508.8	5.4	1829	4	CR599938	CR599938 full-leng
34	508.8	5.4	1829	4	CR601229	CR601229 full-leng
35	508.8	5.4	1829	4	CR617650	CR617650 full-leng
36	508.8	5.4	1829	4	CR622375	CR622375 full-leng
37	508.8	5.4	1834	4	CR597477	CR597477 full-leng
38	508.8	5.4	1837	4	CR626480	CR626480 full-leng
39	508.8	5.4	1839	4	CR604077	CR604077 full-leng
40	508.8	5.4	1842	4	CR597489	CR597489 full-leng
41	508.8	5.4	1861	4	BC028335	BC028335 Homo sapi
42	508.2	5.3	1681	4	CNS0FGAK	CR670454 Tetraodon
43	507.6	5.3	1559	4	CR625663	CR625663 full-leng
44	505.2	5.3	1682	4	CNSOFYTV	CR694503 Tetraodon
45	500.4	5.3	1672	4	CNSOFNQU	CR680138 Tetraodon

OM nucleic - nucleic search, using sw model

March 15, 2006, 22:49:07; Search time 4713 Seconds Run on:

(without alignments)

10999.624 Million cell updates/sec

Title: US-10-694-779-9

Perfect score: 912

1 atgtgcgacggagtccgtag.....cacttcacgcagactcgcac 912 Sequence:

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 5883141 seqs, 28421725653 residues

Total number of hits satisfying chosen parameters: 11766282

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : GenEmbl:*

1: gb_ba:*

2: gb_in:*

3: gb_env:*

4: gb_om:*
5: gb_ov:*

6: gb_pat:*

7: gb_ph:* 8: gb_pr:*

9: gb_ro:*

10: gb_sts:*

11: gb_sy:*

12: gb_un:*

13: gb_vi:*

14: gb_htg:*

15: gb_pl:*

g,

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			₹				
Resi	ılt		Query				
1	No.	Score	Match	Length	DB	ID	Description
	1	912	100.0	912	6	BD133255	BD133255 FOF1-ATPa
	2	912	100.0	9500	6	BD133267	BD133267 FOF1-ATPa
	3	504.2	55.3	8394	1	AB046112	AB046112 Corynebac
	4	504.2	55.3	110000	1	BA000036_12	Continuation (13 o
	5	504.2	55.3	349459	1	BX927151	BX927151 Corynebac
	6	504.2	55.3	349980	6	AX127147	AX127147 Sequence
	7	502.6	55.1	8001	1	AB048368	AB048368 Brevibact
	8	499.2	54.7	110000	1	BA000035 13	Continuation (14 o
	9	488.2	53.5	933	6	AX065629	AX065629 Sequence
	10	456.2	50.0	862	6	AX065631	AX065631 Sequence
	11	454	49.8	347625	1	BX248356	BX248356 Corynebac
	12	443.6	48.6	810	6	BD163537	BD163537 Novel pol
	13	443.6	48.6	810	6	AX121420	AX121420 Sequence
С	14	435	47.7	110000	1	CR931997 15	Continuation (16 o
	15	288.6	31.6	110000	1	AP006618 11	Continuation (12 o
	16	190.8	20.9	110000	1	AE016822 06	Continuation (7 of
	17	190.8	20.9	110000	1	AE016822 07	Continuation (8 of
						-	

	18	134.6	14.8 299050	1	BX251411	BX251411 Tropherym
C	19	134.6	14.8 302529	1	AE016851	AE016851 Tropherym
	20	134.2	14.7 8560	1	SLATPSYNA	Z22606 S.lividans
	21	134.2	14.7 300100	1	SCO939123	AL939123 Streptomy
	22	127.4	14.0 7760	1	AY702091	AY702091 Nonomurae
C	23	120.6	13.2 8323	1	AY488175	AY488175 Bifidobac
С	24	119.4	13.1 110000	1	BA000030_35	Continuation (36 o
C	25	118.8	13.0 110000	1	AE014295_12	Continuation (13 o
C	26	118.8	13.0 349980	6	AX492782	AX492782 Sequence
C	27	118.8	13.0 349980	6	AX553949	AX553949 Sequence
С	28	102	11.2 110000	1	CP000088_28	Continuation (29 o
	- 29	95	10.4 8358	1	AY488174	AY488174 Bifidobac
	30	71.2	7.8 110000	1	BA000012_61	Continuation (62 o
	31	60	6.6 7218	6	I66494	I66494 Sequence 14
	32	59.2	6.5 2220	1	AF054609	AF054609 Brucella
С	33	59.2	6.5 12028	1	AE009590	AE009590 Brucella
	34	59.2	6.5 110000	1	AE014291_03	Continuation (4 of
	35		6.5 110000		AE017223_04	Continuation (5 of
C	36	55.6	6.1 11283		AE005710	AE005710 Caulobact
	37	51.6	5.7 31495		BJU33883	U33883 Bradyrhizob
С	38	51.6	5.7 110000	1	-	Continuation (13 o
C	39	51.6	5.7 110000	1	BA000040_13	Continuation (14 o
	40	50.8	5.6 843	6	AR621598	AR621598 Sequence
С	41	50.8	5.6 1394	6	AR618970	AR618970 Sequence
	42	48.8	5.4 4364			CQ363872 Sequence
C	43	48.8	5.4 110000		AE017283_13	Continuation (14 o
	44	48.4	5.3 12152	1	AE009040	AE009040 Agrobacte
	45	48.4	5.3 12164	1	AE008006	AE008006 Agrobacte

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OM nucleic - nucleic search, using sw model

Run on: March 15, 2006, 05:51:17; Search time 370 Seconds

(without alignments)

4381.449 Million cell updates/sec

Title: US-10-694-779-9

Perfect score: 912

Sequence: 1 atgtgcgacggagtccgtag.....cacttcacgcagactcgcac 912

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 1303057 seqs, 888780828 residues

Total number of hits satisfying chosen parameters: 2606114

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued Patents NA:*

1: /cgn2 6/ptodata/1/ina/1 COMB.seq:*

2: /cgn2_6/ptodata/1/ina/5_COMB.seq:*

3: /cgn2 6/ptodata/1/ina/6A COMB.seq:*

4: /cgn2_6/ptodata/1/ina/6B_COMB.seq:*

5: /cgn2_6/ptodata/1/ina/H_COMB.seq:*

6: /cgn2_6/ptodata/1/ina/PCTUS_COMB.seq:*

7: /cgn2 6/ptodata/1/ina/PP COMB.seq:*

8: /cgn2_6/ptodata/1/ina/RE COMB.seq:*

9: /cgn2_6/ptodata/1/ina/backfiles1.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			₩				
Res	sult		Query				
	No.	Score	Match	Length	DB	ID	Description
-	1	60	6.6	7218	2	US-08-232-463-14	Sequence 14, Appl
	2	50.8	5.6	843	3	US-09-902-540-2999	Sequence 2999, Ap
С	3	50.8	5.6	1394	3	US-09-902-540-365	Sequence 365, App
	4	44.6	4.9	1703	3	US-09-370-253-9	Sequence 9, Appli
С	5	40	4.4	680	3	US-09-533-559-1226	Sequence 1226, Ap
С	6	39	4.3	1581	3	US-09-936-145-15	Sequence 15, Appl
	7	37.8	4.1	702	3	US-09-902-540-7194	Sequence 7194, Ap
C	8	37.8	4.1	6430	3	US-09-902-540-669	Sequence 669, App
	9	37	4.1	663	3	US-09-489-039A-353	Sequence 353, App

	10	36.6	4.0	1865	3	US-09-370-253-5	Sequence 5, Appli
	11	35.6	3.9	1656	3	US-09-902-540-6804	Sequence 6804, Ap
C	12	35.6	3.9	3487	3	US-09-902-540-561	Sequence 561, App
С	13	35	3.8	1083	3	US-09-252-991A-1074	Sequence 1074, Ap
	14	35	3.8	1372	3	US-09-270-767-13796	Sequence 13796, A
	15	35	3.8	1692	3	US-09-252-991A-978	Sequence 978, App
	16	35	3.8	1836	3	US-09-252-991A-1026	Sequence 1026, Ap
	17	34.8	3.8	747	3	US-09-252-991A-14108	Sequence 14108, A
	18	34.8	3.8	1035	3	US-09-252-991A-14017	Sequence 14017, A
С	19	34.8	3.8	1365	3	US-09-252-991A-14122	Sequence 14122, A
	20	34.8	3.8	1574	3	US-09-902-540-2466	Sequence 2466, Ap
	21	34.8	3.8	2019	3	US-09-063-950-3	Sequence 3, Appli
	22	34.8	3.8	2555	3	US-09-866-028-68	Sequence 68, Appl
	23	34.8	3.8	2555	3	US-09-944-457-68	Sequence 68, Appl
	24	34.8	3.8	2555	3	US-09-945-584-68	Sequence 68, Appl
	25	34.8	3.8	2555	3	US-09-944-944-68	Sequence 68, Appl
	26	34.8	3.8	2555	3	US-09-945-587-68	Sequence 68, Appl
	27	34.8	3.8	2768	3	US-09-991-181-51	Sequence 51, Appl
	28	34.8	3.8	2768	3	US-09-990-444-51	Sequence 51, Appl
	29	34.8	3.8	2768	3	US-09-997-333-51	Sequence 51, Appl
	30	34.8	3.8	2768	3	US-09-992-598-51	Sequence 51, Appl
	31	34.8	3.8	2852	3	US-09-063-950-1	Sequence 1, Appli
С	32	34.8	3.8	14077	3	US-09-902-540-1109	Sequence 1109, Ap
	33	34.4	3.8	505	3	US-09-621-976-15639	Sequence 15639, A
	34	34.4	3.8	219964	3	US-09-949-016-15086	Sequence 15086, A
	35	34	3.7	1080	3	US-09-875-076-27	Sequence 27, Appl
	36	34	3.7	3626	3	US-09-620-312D-93	Sequence 93, Appl
С	37	33.8	3.7	14807	3	US-09-902-540-1085	Sequence 1085, Ap
С	38	33.6	3.7	525	3	US-09-252-991A-9910	Sequence 9910, Ap
С	39	33.6	3.7	723	3	US-09-252-991A-10004	Sequence 10004, A
	40	33.6	3.7	744	3	US-09-252-991A-10165	Sequence 10165, A
	41	33.6	3.7	870	3	US-09-252-991A-6993	Sequence 6993, Ap
	42	33.6	3.7	1191	3	US-09-252-991A-7232	Sequence 7232, Ap
С	43	33.6	3.7	1218	3	US-09-252-991A-7465	Sequence 7465, Ap
	44	33.6	3.7	1248	3	US-09-252-991A-7180	Sequence 7180, Ap
	45	33.6	3.7	1254	3	US-09-252-991A-10100	Sequence 10100, A
							<u>-</u>

OM nucleic - nucleic search, using sw model

Run on: March 16, 2006, 15:33:28; Search time 434 Seconds

(without alignments)

4899.806 Million cell updates/sec

Title: US-10-694-779-9

Perfect score: 912

Sequence: 1 atgtgcgacggagtccgtag.....cacttcacgcagactcgcac 912

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 8023312 seqs, 1165852854 residues

Total number of hits satisfying chosen parameters: 16046624

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications NA New:*

- /cgn2 6/ptodata/2/pubpna/US08 NEW PUB.seq:*
- /cgn2_6/ptodata/2/pubpna/US06_NEW_PUB.seq:*
- /cgn2_6/ptodata/2/pubpna/US07_NEW_PUB.seq:*
- /cgn2_6/ptodata/2/pubpna/PCT_NEW_PUB.seq:*
- /cgn2_6/ptodata/2/pubpna/US09 NEW PUB.seq:*
- /cgn2_6/ptodata/2/pubpna/US09 NEW PUB.seq1:*
- /cgn2_6/ptodata/2/pubpna/US10_NEW_PUB.seq:* 7:
- 8: /cgn2_6/ptodata/2/pubpna/US10_NEW_PUB.seq1:* /cgn2_6/ptodata/2/pubpna/US11_NEW_PUB.seq:*
- 10: /cgn2_6/ptodata/2/pubpna/US11_NEW_PUB.seq2:*
- 11: /cgn2 6/ptodata/2/pubpna/US11 NEW PUB.seq3:*
- 12: /cgn2 6/ptodata/2/pubpna/US11 NEW PUB.seq4:*
- 13: /cgn2 6/ptodata/2/pubpna/US60 NEW PUB.seq:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	Query Match	Length	DB	ID	Description
1	36.4	4.0	1536	9	US-11-096-568A-22147	Sequence 22147, A
2	36.4	4.0	1791	9	US-11-096-568A-18969	Sequence 18969, A
3	36.2	4.0	1792	9	US-11-096-568A-23390	Sequence 23390, A
4	35.8	3.9	1400	12	US-11-136-527-7196	Sequence 7196, Ap

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                         1745
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                                                                Sequence 5841, Ap
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